

Evolutionary Relationships among Parvoviruses: Virus-Host Coevolution among Autonomous Primate Parvoviruses and Links between Adeno-Associated and Avian Parvoviruses

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The current classification of parvoviruses is based on virus host range and helper virus dependence, while little data on evolutionary relationships among viruses are available. We identified and analyzed 472 sequences of parvoviruses, among which there were (virtually) full-length genomes of all 41 viruses currently recognized as individual species within the family *Parvoviridae*. Our phylogenetic analysis of full-length genomes as well as open reading frames distinguished three evolutionary groups of parvoviruses from vertebrates: (i) the human helper-dependent adeno-associated virus (AAV) serotypes 1 to 6 and the autonomous avian parvoviruses; (ii) the bovine, chipmunk, and autonomous primate parvoviruses, including human viruses B19 and V9; and (iii) the parvoviruses from rodents (except for chipmunks), carnivores, and pigs. Each of these three evolutionary groups could be further subdivided, reflecting both virus-host coevolution and multiple cross-species transmissions in the evolutionary history of parvoviruses. No parvoviruses from invertebrates clustered with vertebrate parvoviruses. Our analysis provided evidence for negative selection among parvoviruses, the independent evolution of their genes, and recombination among parvoviruses from rodents. The topology of the phylogenetic tree of autonomous human and simian parvoviruses matched exactly the topology of the primate family tree, as based on the analysis of primate mitochondrial DNA. Viruses belonging to the AAV group were not evolutionarily linked to other primate parvoviruses but were linked to the parvoviruses of birds. The two lineages of human parvoviruses may have resulted from independent ancient zoonotic infections. Our results provide an argument for reclassification of *Parvovirinae* based on evolutionary relationships among viruses.

The virus family *Parvoviridae* comprises small animal viruses with linear single-stranded DNA genomes. The genomes of parvoviruses are about 5 kb in length and contain two large open reading frames (ORFs). The first codes for two nonstructural proteins, NS-1 and NS-2, while the second encodes coat proteins VP-1 to VP-3 (or two of them), which have substantial amino acid identity, being derived from overlapping reading frames (for a review, see reference 12).

As now classified, the family *Parvoviridae* contains two subfamilies: the *Parvovirinae*, or viruses from vertebrates, and the *Densovirinae*, or viruses from insects and (tentatively) other arthropods (62). The subfamily *Parvovirinae* contains three genera: *Parvovirus*, comprising most parvoviruses from vertebrates; *Erythrovirus*, comprising B19 and V9 parvoviruses as well as parvoviruses from rhesus and pig-tailed macaques, and *Dependovirus*, which comprises adeno-associated viruses (AAV). The last two genera include human viruses: the B19 and V9 parvoviruses (*Erythrovirus*) and AAV serotypes 1 to 6 (*Dependovirus*). Within *Densovirinae*, four genera are recognized. The current classification of parvoviruses is based primarily on their host range and their dependence on help from other viruses for replication, according to the traditional separation of parvoviruses into three types: (i) autonomous viruses of vertebrates, (ii) helper-dependent viruses of vertebrates, and (iii) autonomous viruses of insects (62).

The relationships among parvoviruses have been extensively studied by using serological methods as well as DNA hybridization and restriction mapping analyses. A relatively high sequence homology of goose and Muscovy duck autonomous parvoviruses (GPV and MDPV, respectively) with helper-dependent AAV-2, but not with other autonomous parvoviruses, has been documented (18, 68). Although direct data on sequence homology of GPV and MDPV to AAV serotypes other than AAV-2 were not available, DNA cross hybridization data have suggested that GPV is even more similar to AAV-1 and AAV-3 (18). On the other hand, little similarity between the two groups of human parvoviruses, B19 and AAV, has been observed (68). Feline panleukopenia virus, canine parvovirus, and mink enteritis virus (MEV) are highly homologous and classified as host range variants of the feline parvovirus (FelinePV) (reviewed in reference 46), but another mink parvovirus, the Aleutian mink disease virus (AMDV), has little homology with MEV (14). These observations prompted the suggestion that the original hypothesis of a host-dependent evolution of parvoviruses (8) may have limited value both within and among genera (49, 68). Furthermore, autonomous parvoviruses are dependent on helper functions that are transiently expressed in host cells, and helper viruses can substantially increase their replication, while helper-dependent viruses can replicate autonomously under certain conditions (12). These observations together with genetic homology between some autonomous and helper-dependent viruses resulted in the validity of the other main criterion for classification of parvoviruses, dependence on helper viruses, also being ques-

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tioned (18, 68). The distinction that autonomous parvoviruses encapsidate primarily DNA strands that are complementary to mRNA, whereas AAV encapsidate strands of either polarity with equal frequency, is also far from absolute. For instance, bovine parvovirus encapsidates up to 30% of DNA strands with the same polarity as that of mRNA. In certain hosts, the autonomous parvovirus LuIII encapsidates strands with either polarity in equal measure (12).

Over the last decade, a massive amount of genetic information has been obtained for various virus groups. For several groups, including the human immunodeficiency viruses (HIV) and hepatitis C viruses, genetic classifications that reflect evolutionary relationships have been developed (38, 39, 57). However, no systematic and explicit study on evolutionary relationships among *Parvoviridae* has yet been performed, although (virtually) full-length genomes of several members of each of the recognized genera are available. Such a study is essential to elucidate the principal issues of parvovirus biology, including the evolutionary relationships both among and within subfamilies and genera, the driving forces of parvovirus evolution, and possible cross-species transmissions. In the present study, we address these basic issues and analyze currently available sequence information on parvoviruses by using phylogenetic methods.

MATERIALS AND METHODS

Sequences. In the GenBank, we identified 472 sequences of parvoviruses for use in this study. They were retrieved by using Batch Entrez software, which allows a search for sequences belonging to a specified organism (<http://www.ncbi.nlm.nih.gov/Entrez/batch.html>). We specified *Parvoviridae* as the organism name, according to the taxonomy database at the National Center for Biotechnology Information, and performed an additional search for *Parvovirus*. We used the classification of parvoviruses accepted by the International Committee on Taxonomy of Viruses (<http://www.ncbi.nlm.nih.gov/htbin-post/Taxonomy/>). The viruses we studied are referred to by their descriptive name (e.g., FelinePV) or trivial name (e.g., B19), as it is used in the nomenclature of parvoviruses (62). The sequences are referred to by their GenBank accession numbers, and the reference information is provided in Table 1.

Sequence analysis. The BioEdit, version 4.8.6, software (28) was used to manipulate the retrieved sequences. The alignment of sequences was performed by using the ClustalW software (60). For full-length genomes as well as noncoding regions, nucleotide sequences were aligned. For coding regions, the alignment was performed for amino acid sequences.

Phylogenetic analysis was performed by using several methods. For all methods, positions containing an alignment gap were excluded from pairwise sequence comparisons. Bootstrap resampling was performed for each analysis (100 replications). Nucleotide distances were analyzed by using the neighbor-joining algorithm as implemented in the PHYLIP package (NEIGHBOR), based on the Kimura two-parameter distance estimation method or the proportion of differences (p distance). For coding regions, additional analyses of nonsynonymous and synonymous nucleotide substitutions (those which change or do not change the amino acid, respectively) was performed by using the MEGA software (37). Estimation of both synonymous distances (Ds) and nonsynonymous distances (Da) was based on the Nei-Gojobori method (37). The ratios of synonymous to nonsynonymous substitutions (Ds/Da) were calculated (41).

Recombination analysis was performed by using the bootscanning method as implemented in the SimPlot software (available at <http://www.med.jhu.edu/deptmed/sray/>).

Many viruses are represented in the GenBank by single full-length genome sequences, but more than one sequence are available for several viruses. For B19 virus, we used full-length sequences but also about 200 shorter sequences, typically a few hundred nucleotides in length. These partial genomes were aligned with all full-length genome sequences, and the B19 consensus sequence was calculated as the arithmetic mean of all nucleotides or amino acids at a particular position (39, 42). This consensus sequence was used in the analyses together with the individual full-length genomes.

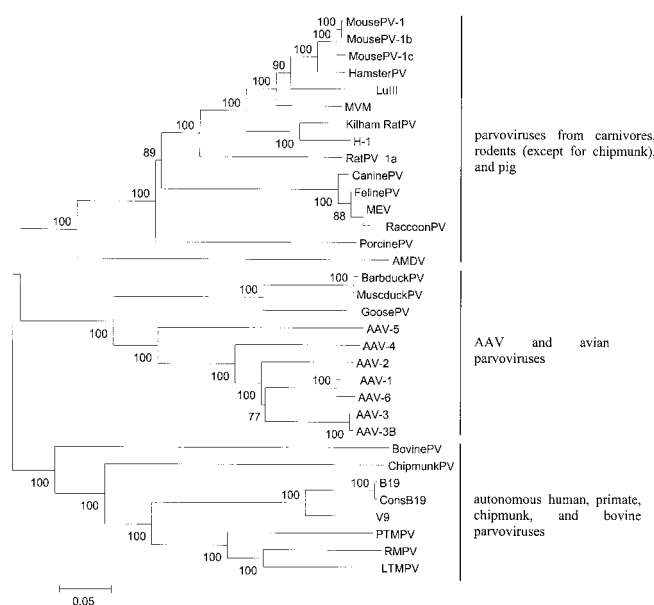


FIG. 1. The three evolutionary groups of *Parvovirinae*. The neighbor-joining phylogenetic tree is based on the analysis of (virtually) full-length genomes of all members of the *Parvovirinae* subfamily that are recognized as individual virus species, one sequence per species (except for the B19 virus, for which a consensus, ConsB19, of 215 available sequences is also included). For RaccoonPV, only a shorter sequence is available. Bootstrap values are shown (100 replications). Sequences used in this analysis are in boldface in Table 1. For virus abbreviations, see Table 1.

RESULTS

Identification of phylogenetic groups within *Parvovirinae*.

To identify groups of phylogenetically related viruses within *Parvovirinae*, we analyzed (virtually) full-length genomes of all members of the three genera that are distinguished by the International Committee on Taxonomy of Viruses as distinct virus species. *Parvovirus* species included were bovine, simian (from the cynomolgus [long-tailed] macaque), Manchurian chipmunk, canine, feline panleukopenia, Georgian raccoon (only a partial sequence of 2,410 nucleotides in length is available), porcine, mice minute, mouse 1, mouse 1b, mouse 1c, rat 1a, Kilham rat, hamster, LuIII, Barbarie duck, and H1 parvoviruses as well as MEV, AMDV, GPV, and MDPV. *Erythrovirus* species included an individual B19 virus and the consensus of 215 B19 sequences and V9 and rhesus and pig-tailed macaque parvoviruses. *Dependovirus* species included AAV serotypes 1 to 6. The list of sequences analyzed is provided in Table 1.

In total, genomic sequences of 32 virus species were aligned. Based on phylogenetic analysis, they fell into three groups (Fig. 1): (i) AAV serotypes 1 to 6 and GPV, Barbarie duck parvovirus, and MDPV; (ii) primate (B19, V9, and three viruses from macaques), chipmunk, and bovine parvoviruses; (iii) parvoviruses from all rodents (except for chipmunks), carnivores, and pigs.

Additionally, we analyzed viruses from the *Densovirinae* subfamily (Table 1). None of viruses from the *Densovirinae* clustered together with *Parvovirinae* (data not shown).

AAV and avian parvoviruses. To analyze phylogenetic relationships among AAV and avian parvoviruses, we aligned sequences of viruses belonging to this phylogenetic group. The

TABLE 1. Virus sequences used in this study^a

Virus species	Abbreviated name(s)	Accession no.	Reference ^b
<i>Parvovirinae</i>			
<i>Parvovirus</i>			
Bovine parvovirus	BovinePV, BPV	M14363	20
Canine parvovirus	CaninePV, CPV	M19296	51
		M38245	47
		D26079	34
Minute virus of mice	MVM	J02275	3
		X02481	54
		V01115	3
		M12032	2
		U34256	13
Mouse parvovirus 1	MousePV-1	U12469	5
Mouse parvovirus 1b	MousePV-1b	U34253	13
Mouse parvovirus 1c	MousePV-1c	U34254	13
LuIII		M81888	24
Barbarie duck parvovirus	BarbduckPV, BDPV	U22967	68
Feline panleukopenia virus	FelinePV, FPV, FPLV	M38246	47
		X55115	44
		M10824	19
Goose parvovirus	GoosePV, GPV	U25749	68
Mink enteritis virus	MEV	D00765	36
Porcine parvovirus	PorcinePV, PPV	L23427	11
		M38367	64
		D00623	50
		M32787(orf2)	63
		U44978	10
Rat parvovirus 1a	RatPV-1a	AF036710	6
Kilham rat virus	KilhamRatPV	U79033	Brown and Like, unpub., 1996
		AF036711	6
Aleutian mink disease parvovirus	AMDV	M20036	14
		X97629	56
		Z18276	Perryman et al., unpub., 1992
Hamster parvovirus	HamsterPV	U34255	13
Muscovy duck parvovirus	MuscduckPV, MDPV	X75093	68
Raccoon parvovirus, Georgia raccoon virus	RaccoonPV	M24005 (partial)	48
Parvovirus H1	H1	X01457	52
Simian parvovirus from cynomolgus monkeys (long-tailed macaque)	LTMPV	U26342	17
Chipmunk parvovirus	ChipmunkPV	U86868	67
<i>Erythrovirus</i>			
B19 virus		AF162273	Gallinella and Venturoli, unpub., 1999
		AF161226	33
		AF161225	33
		AF161224	33
		AF161223	33
		AF113323	31
		AB030694	Ishii et al., unpub., 1999
		AB030693	Ishii et al., unpub., 1999
		AB030673	61
		Z70599	32
		Z70560	32
		M24682	15
V9 virus		AX003421	4
Rhesus macaque parvovirus	RMPV	AF221122	27
Pig-tailed macaque parvovirus	PTMPV	AF221123	27
<i>Dependovirus</i>			
Adeno-associated virus serotype 1	AAV-1	AF063497	65
Adeno-associated virus serotype 2	AAV-2	J01901	55
Adeno-associated virus serotype 3	AAV-3	U48704	45
Adeno-associated virus serotype 3B	AAV-3b	AF028705	53
Adeno-associated virus serotype 4	AAV-4	U89790	22
Adeno-associated virus serotype 5	AAV-5	AF085716	21
Adeno-associated virus serotype 6	AAV-6	AF028704	53
<i>Densovirinae</i>			
<i>Junonia coenia</i> densovirus	JcDNV	S47266	25
<i>Galleria mellonella</i> densovirus	GmDNV	L32896	58
<i>Bombyx mori</i> densovirus	BmDNV	AB042597	Nonaka et al., unpub., 2000
<i>Aedes albopictus</i> parvovirus	AaPV	X74945	16
<i>Aedes aegypti</i> densovirus	AedesDNV	M37899	1
<i>Periplaneta filiginosa</i> densovirus	PfDNV	AF192260	Guo et al., unpub., 1999
		AB028936	66
<i>Diatraea saccharalis</i> densovirus	DsDNV	AF036333	Boublik et al., unpub., 1997
Yamanashi isolate from silkworm		S78547	7
Parvovirus-like virus		AB033596	29
Infectious hypodermal and hematopoietic necrosis virus from penaeid shrimps		AF218266	43

^a Sequences in boldface are used in Fig. 1.^b Unpub., unpublished data.

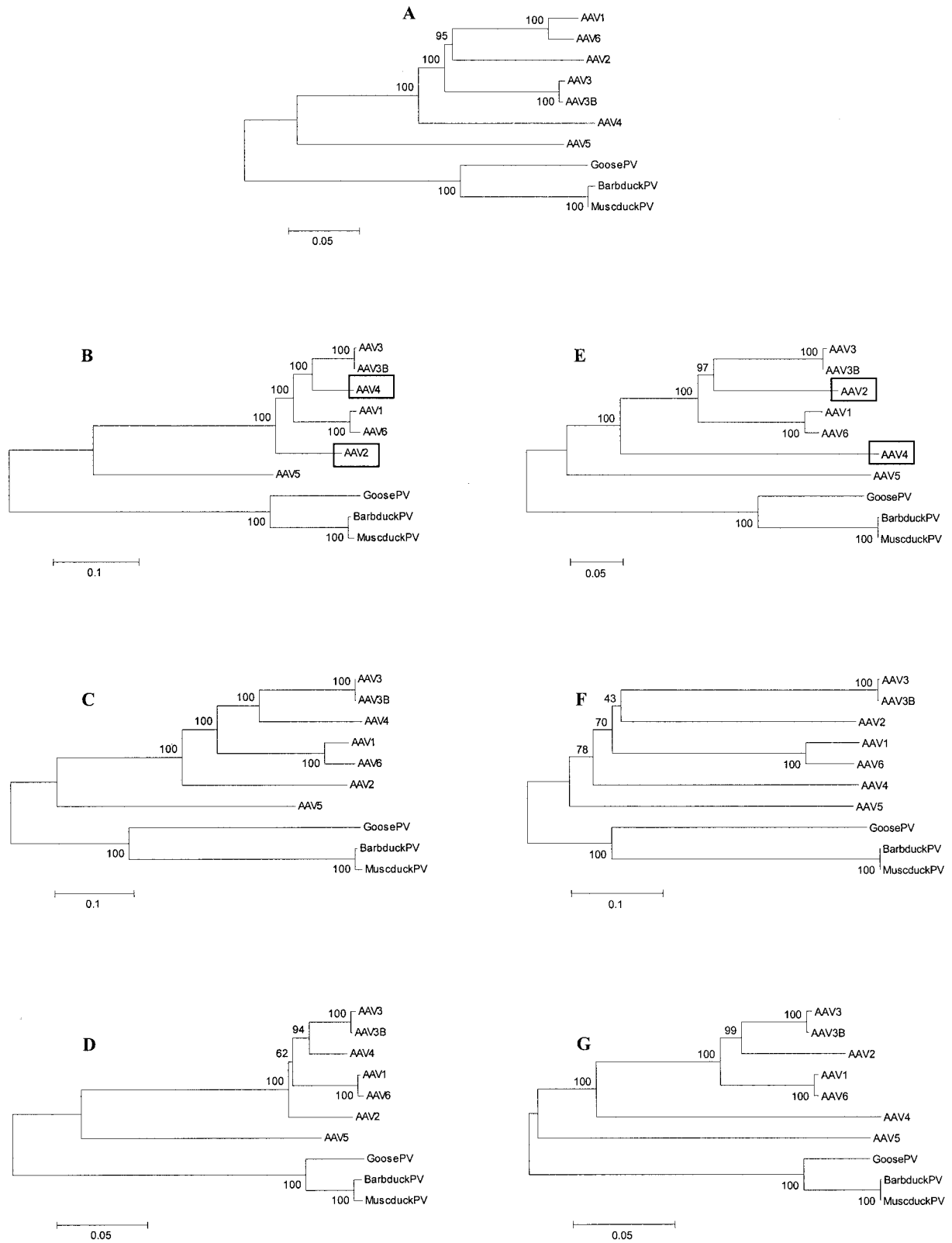


FIG. 2. Phylogenetic relationships among the AAV serotypes 1 to 6 and parvoviruses from GPV, Barbarie duck parvovirus (BarbduckPV), and MDPV (MuscduckPV). Bootstrap values are shown (100 replications). (A) Relationships based on nucleotide p distances among full-length genome sequences; (B to D) relationships based on nucleotide Kimura two-parameter distances, D_s and D_a , respectively, for orf1; (E to G) nucleotide distances, D_s and D_a , respectively, for orf2. For panels B and E, positions of AAV-2 and AAV-4 are marked. Virus abbreviations are listed in Table 1.

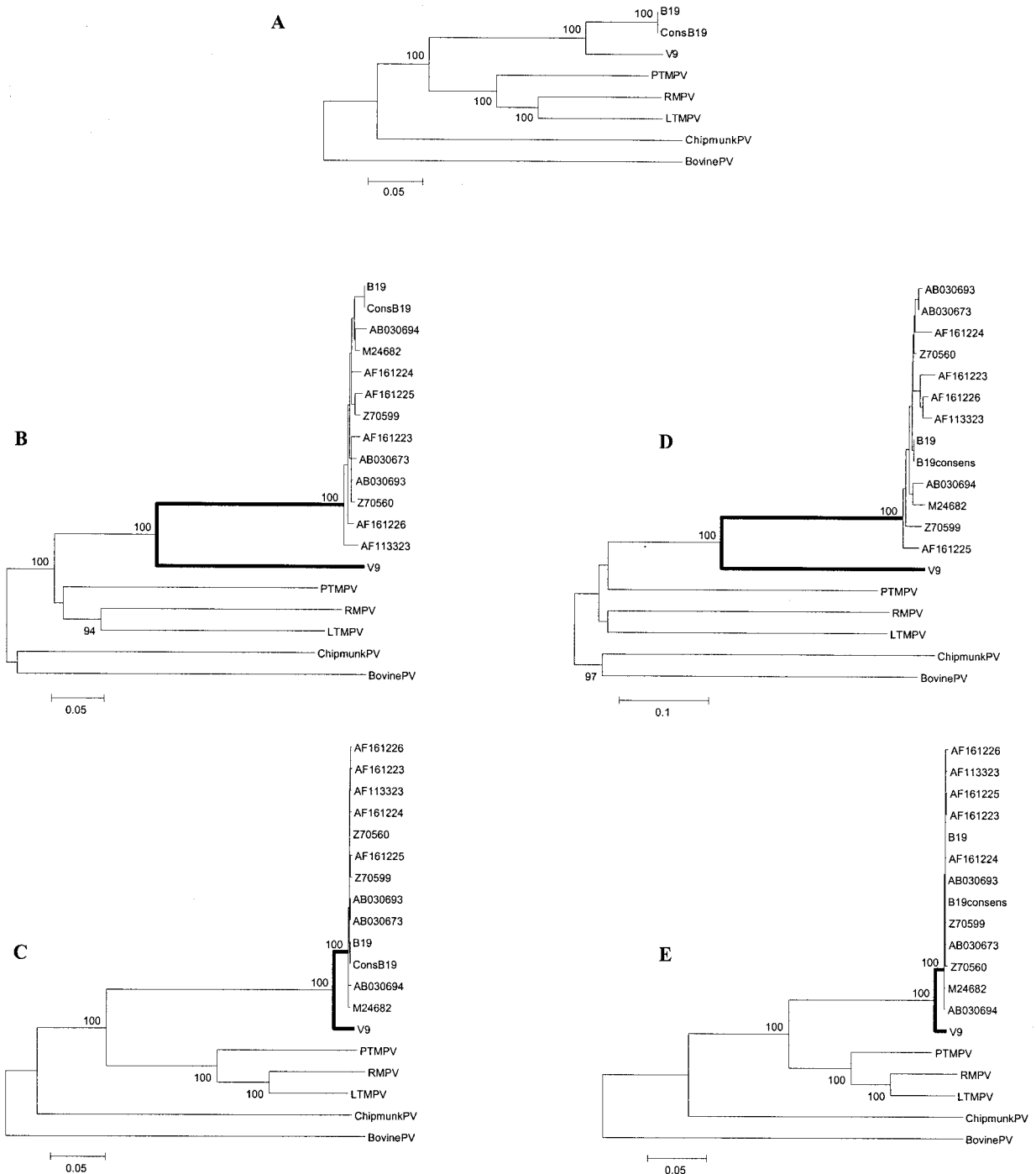


FIG. 3. Phylogenetic relationships among the autonomous primate, chipmunk, and bovine parvoviruses. In addition to sequences used in Fig. 1, 11 more sequences of B19 are included (labeled by their GenBank accession numbers). Bootstrap values above 70 are shown (100 replications). (A) Relationships based on nucleotide p distances among full-length genome sequences; (B and C) relationships based on Ds and Da, respectively, for orf1; (D and E) relationships based on Ds and Da, respectively, for orf2. Branches between the B19 cluster and V9 are in boldface (B to E). Virus abbreviations are in Table 1.

analyses were based on nucleotide distances as well as, for coding regions, Ds and Da.

Irrespective of the phylogenetic model and genomic region used, the three avian parvoviruses clustered together and separately from AAV, with a bootstrap value of 100 (Fig. 2). The two viruses from ducks were virtually identical, with their Ds

and Da being 0.01 for orf1 and 0.00 for orf2. Among AAV, two pairs of closely related viruses were found. Besides the two sequences belonging to viruses from the same serotype (AAV-3 and AAV-3B), AAV-1 and AAV-6 also clustered together. Although these two viruses are defined as separate AAV serotypes, the vast majority of nucleotide substitutions

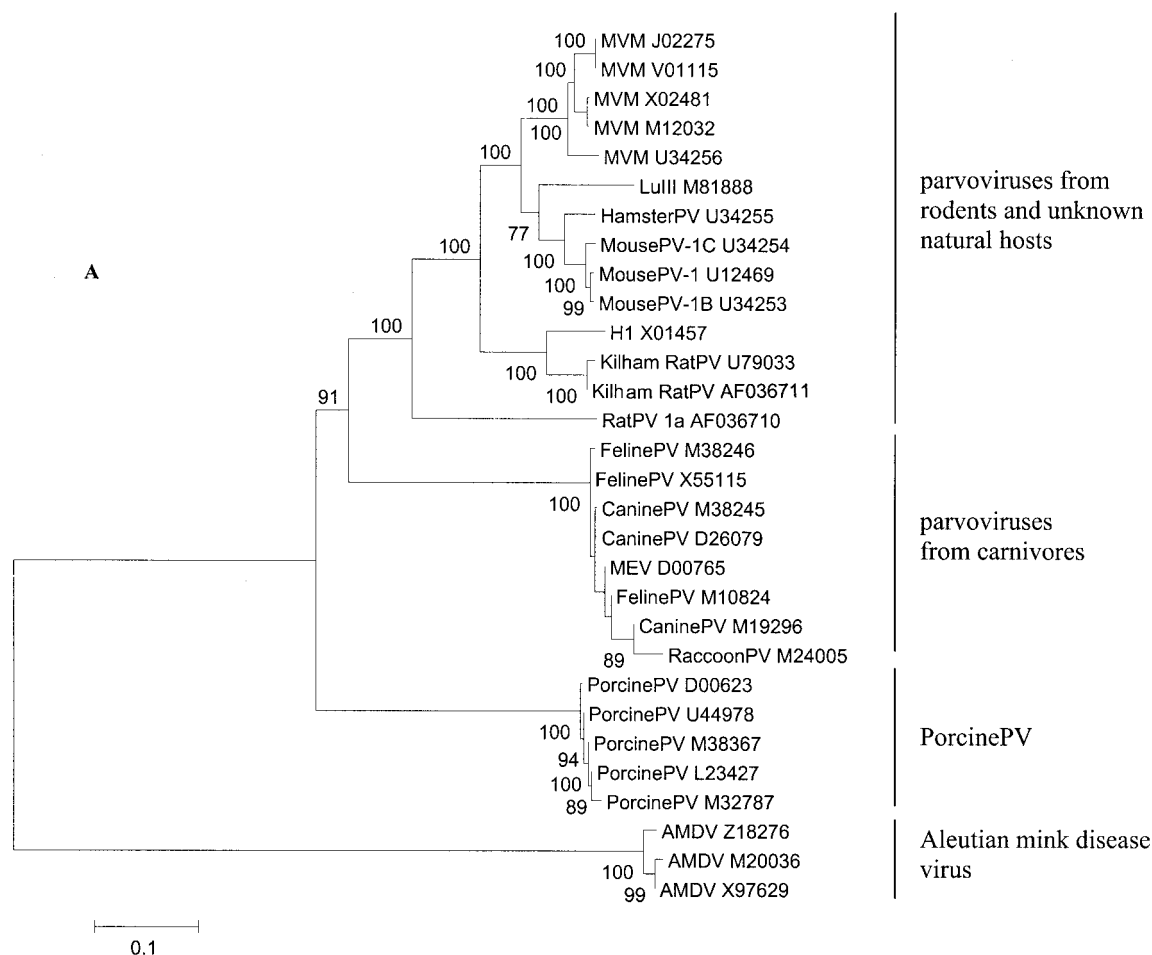


FIG. 4. Phylogenetic relationships among parvoviruses from rodents, carnivores, and pigs. The four phylogenetic subgroups are shown. (A) All full-length sequences available for each virus species are included (labeled by their virus names and the GenBank accession numbers). Bootstrap values above 70 are shown (100 replications). (A) relationships based on nucleotide p distances among full-length genome sequences; (B to D) relationships based on the nucleotide Kimura two-parameter distances, D_s , and D_a , respectively, for orf1; (E to G) relationships based on nucleotide Kimura two-parameter distances, D_s , and D_a , respectively, for orf2. In orf1 (B to D), MousePV, HamsterPV, and MVM form a homogeneous cluster (B, grey box), to which LuIII (arrow) is an outlier. In contrast, in orf2 (E to G), MousePV and HamsterPV (E, open box) cluster with LuIII (arrow) and not with MVM (grey box). Virus abbreviations are in Table 1.

between AAV-1 and AAV-6 are synonymous, with D_s being 0.07 and 0.11 for orf1 and orf2, respectively, and D_a being 0.00 for both ORFs. AAV-3 and AAV-3B, AAV-1 and AAV-6, and AAV-2 were approximately equidistant from each other as well as from AAV-5, which appeared to be the most distantly related to other AAV (Fig. 2). Within this virus group, branching orders of two viruses, AAV-2 and AAV-4, varied with the genetic region analyzed. The orf1 sequence of AAV-4 clustered together with AAV-3 and AAV-3B, AAV-1 and AAV-6, and AAV-2 and was most closely related to AAV-3 (Fig. 2B to D), whereas the orf2 sequence of AAV-4 branched out between AAV-5 and the main cluster of AAV (Fig. 2E to G). The position of AAV-2 (within or outside the AAV-3 and AAV-3B and AAV-1 and AAV-6 clusters) also depended upon the genetic region (Fig. 2).

For all pairwise sequence comparisons, the D_s/D_a ratios were markedly higher than 1. The D_a between any two sequences did not exceed 0.39 (AAV-1 and AAV-6 versus avian parvoviruses; orf1), but the vast majority of pairwise D_s were

higher (Fig. 2D and G versus C and F). Remarkable differences between the D_s and D_a were observed for GPV versus duck parvoviruses, for which the D_s were 0.58 to 0.59, compared to D_a of 0.06 to 0.07. Among AAV, the most remarkable differences between D_s and D_a were observed for the comparisons of AAV-2 and AAV-4 in orf1, 0.44 versus 0.07, and AAV-1 and AAV-6 and AAV-3 and AAV-3B in orf 2, 0.52 to 0.57 versus 0.08 to 0.09 (Fig. 2).

Autonomous primate, chipmunk, and bovine parvoviruses. In addition to a single B19 individual sequence and the B19 consensus, we analyzed all 12 individual B19 sequences for which both orf1 and orf2 regions are available.

For this group of viruses, topologies of phylogenetic trees were virtually identical when based on full-length sequence analysis or D_a in orf1 and orf2 (Fig. 3). B19 and V9 clustered together, as did the simian parvoviruses, whereas chipmunk and bovine parvoviruses (ChipmunkPV and BovinePV) were outliers. Among B19 viruses, high genetic homogeneity was observed. Within the two ORFs, the mean D_s among B19

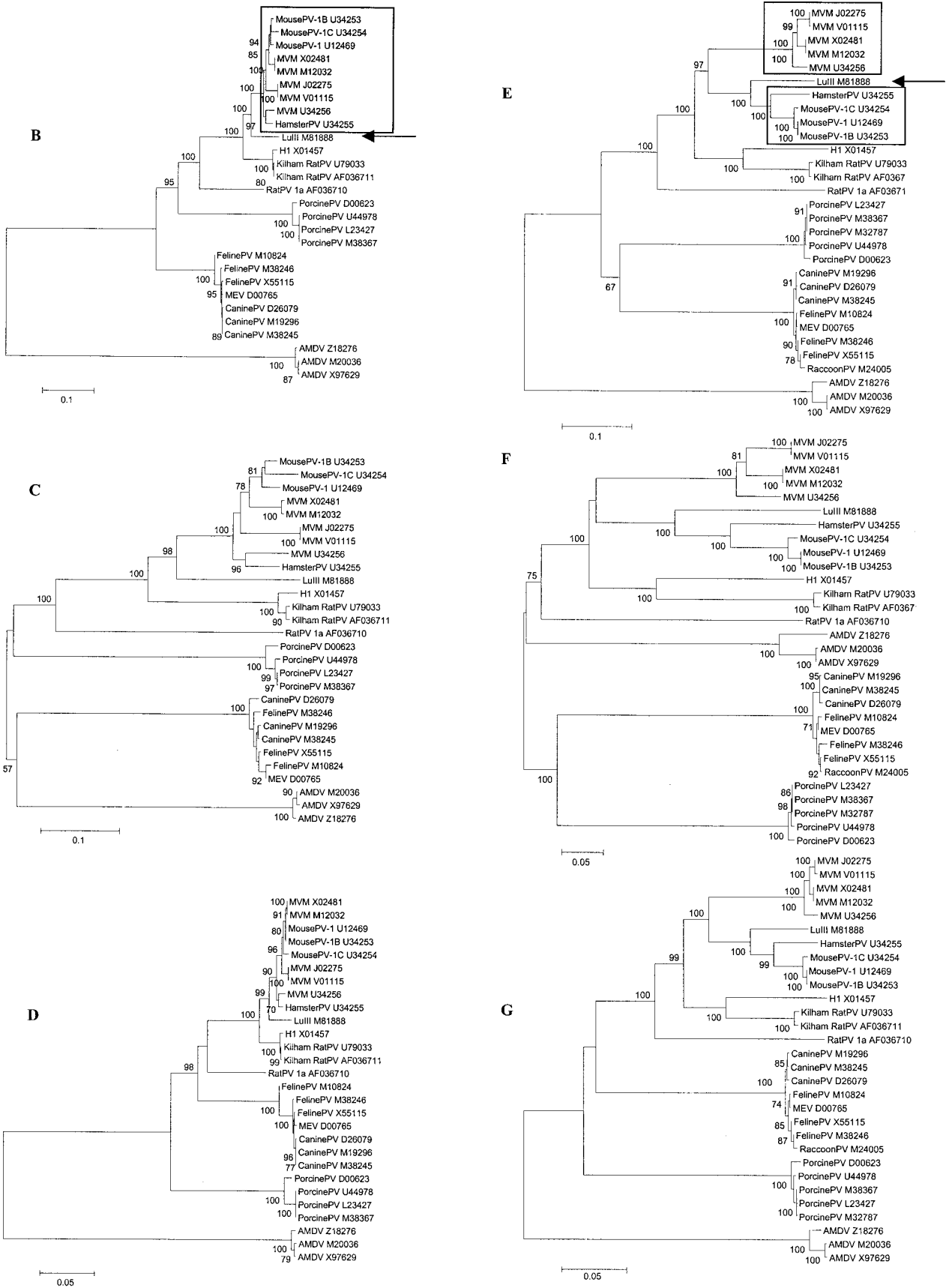


FIG. 4—Continued.

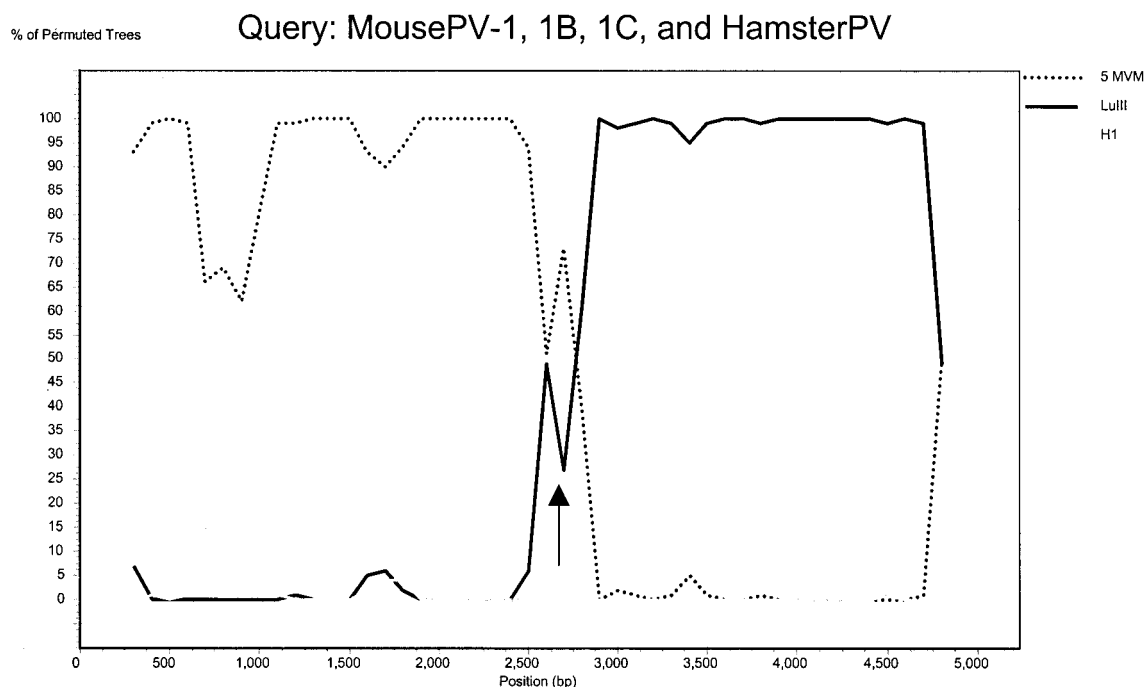


FIG. 5. Bootscan analysis of the phylogenetic relationships among LuIII and parvoviruses from mice and hamsters. The three MousePV and HamsterPV were used as a query sequence group in comparison to the five MVM (comparison group 1), LuIII (comparison group 2), and H1 (outgroup). Analysis settings were as follows: window size, 400 nucleotides; step 100 nucleotides; bootstrap resampling, 100; distance, Kimura two-parameter distance; transitions/transversions ratio, 2. Arrow, recombination site.

viruses were 0.022 (range, 0.007 to 0.034) and 0.035 (range, 0.007 to 0.054) and the mean D_a were 0.003 (range, 0.001 to 0.005) and 0.002 (range, 0.001 to 0.005), resulting in the mean D_s/D_a ratios of 7.3 and 17.5 for orf1 and orf2, respectively. Negative selection was even more evident in our comparison of the two human parvoviruses: while the D_s between the V9 sequence and the B19 consensus were 0.40 and 0.45, the D_a were 0.03 and 0.02 (Fig. 3B to E), resulting in mean D_s/D_a ratios of 13.3 and 22.5 for orf1 and orf2, respectively. Pairwise D_s between human and simian parvoviruses were also markedly higher than D_a (Fig. 3). Moreover, phylogenetic analysis of orf2 based on D_s resulted in virtually complete loss of tree structure as B19 and V9, the three macaque viruses, ChipmunkPV, and BovinePV were equidistant from each other (Fig. 3D).

Parvoviruses from rodents, carnivores, and pigs. For several viruses within this evolutionary group, more than one full-length sequence were available, permitting study of genetic heterogeneity within virus species. We included five strains of minute virus of mice (MVM), two Kilham rat parvoviruses (KilhamRatPV), three FelinePV, three canine parvoviruses (CaninePV), five porcine parvoviruses (PorcinePV), and three AMDV sequences in the analysis.

Our analysis distinguished four major subgroups of evolutionarily related viruses: (i) viruses from rodents and unknown natural hosts, (ii) viruses from carnivores, except for AMDV, (iii) PorcinePV, and (iv) AMDV, which was the most distantly related to all other viruses in this group (Fig. 4A). The four subgroups were observed in all phylogenetic trees (Fig. 4), but their branching order varied. Based on D_a , viruses of rodents, carnivores, and pigs clustered together and were approximately equidistant from each other (Fig. 4D and G), while AMDV ap-

peared to be an outlier. Based on synonymous distances, all four subgroups were equidistant from each other (Fig. 4C and F).

Similar to what was observed for AAV, avian parvoviruses, and primate parvoviruses, the mean D_s/D_a ratios for pairwise comparisons within groups were above 1 (range: 1.3 to 10.3), except for PorcinePV in orf2, a reflection of its extreme genetic homogeneity ($D_s = 0.002$, $D_a = 0.003$). The largest D_s/D_a ratio was observed for the comparisons of mouse and hamster viruses with LuIII in orf1: 0.30/0.04 (Fig. 4C and D).

Among the rodent viruses, we identified three subclusters, which comprised (i) LuIII and viruses from mice and hamsters, (ii) KilhamRatPV and H1, and (iii) the most distantly related rat parvovirus (RatPV) (Fig. 4). The second subgroup of viruses from various natural hosts, viruses from carnivores, and the two subgroups of viruses from single hosts, PorcinePV and AMDV, were much more homogeneous. Among viruses from carnivores, the FelinePV, raccoon parvovirus (RaccoonPV), and MEV clustered together and separately from CaninePV when D_a were analyzed (Fig. 4D and G). This trend was less pronounced for D_s (Fig. 4C and F). Genetic heterogeneity in FelinePV was higher than that in CaninePV. The mean D_s/D_a ratios among FelinePV and CaninePV and between these two viruses were 0.016/0.004, 0.008/0.001, and 0.023/0.004, respectively, for orf1 and 0.016/0.004, 0.004/0.002, and 0.023/0.007, respectively, for orf2. Within these three subgroups, both the mean D_s and D_a were generally below 0.02.

For the three subclusters of rodent parvoviruses, RatPV was an outlier in all phylogenetic trees. For the other two subclusters, remarkable patterns were identified. For the H1-KilhamRatPV subcluster, we observed a great difference in the evolutionary distances between the two viruses for the two ORFs.

Within orf1, the mean Da and Ds between H1 and the two KilhamRatPV were 0.004 and 0.044, respectively, while the Da and Ds within orf2 were 34 and 9 times greater and equal to 0.135 and 0.388, respectively (Fig. 4).

For the mouse-hamster-LuIII subcluster, even more complex relations among viruses were found. In orf1, MVM, mouse parvovirus (MousePV), and hamster parvovirus (HamsterPV) represented an extremely homogeneous (mean Da = 0.01, Ds = 0.11) monophyletic group, to which LuIII was an outlier (Fig. 4B to D; bootstrap value of 100). Within this cluster, sequences belonging to distinct virus species were intermixed (Fig. 4B). In contrast, our analysis of orf2 revealed that MousePV and HamsterPV cluster together with LuIII and are distant (mean Da = 0.18, Ds = 0.56) from MVM (Fig. 4E to G; bootstrap value of 100). In orf2, sequences belonging to all recognized virus species represented monophyletic groups (Fig. 4E to G). Yet no host-related clustering was observed, as sequences of MousePV clustered together with HamsterPV and LuIII and not with sequences of another mouse virus, MVM. The mosaicism of virus genomes within this subcluster was further supported by our analysis of full-length sequences with the bootscanning method (Fig. 5). In this analysis, MousePV-1, -1B, -1C, and HamsterPV (query group) were compared to five full-length sequences of MVM (comparison group 1) and LuIII (comparison group 2), whereas the sequence of H1 was used as an outgroup. While the left part (positions 1 to 2600) of the MousePV and HamsterPV genomes clustered together with MVM, the right part of the genomes clustered together with LuIII and not with MVM (Fig. 5).

DISCUSSION

Currently, the GenBank database contains sequences of about 500 viruses from the *Parvoviridae* family. The vast majority of them are vertebrate viruses, while for viruses of invertebrates only a dozen sequences are available.

So far, no systematic evolutionary study has been performed on parvoviruses. Typically, earlier studies focused on describing the amino acid identities of a new virus with a few of the most closely related sequences (5, 6, 9, 27, 53, 67, 68). To the best of our knowledge, the only parvoviruses for which evolutionary issues have been specifically addressed by analyzing full-length genomes are FelinePV and CaninePV (35). Due to the lack of systematically analyzed data, genetic information was not used as the basis for parvovirus classification.

In the present study, we used all available sequence information and powerful phylogenetic methods to learn whether the *Parvoviridae* are evolutionarily related viruses and whether their current classification into subfamilies and genera truly reflects the evolutionary relationships among viruses. Moreover, we attempted to study how and to what degree various evolutionary factors, such as positive or negative selection, recombinations, host-dependent evolution, cross-species transmissions, and the independent evolution of genomic regions, were operational during the evolution of parvoviruses.

Toward the evolutionary classification of *Parvovirinae*. Our analysis of the genomes of 32 parvoviruses, all recognized virus species for which (virtually) full-length genome sequences are available, revealed the existence of three groups of evolutionarily related viruses (Fig. 1 to 4): (i) AAV and all three known

avian parvoviruses; (ii) all five known autonomous primate parvoviruses, ChipmunkPV, and the outlier BovinePV; (iii) parvoviruses from rodents (except for chipmunks), carnivores, and pigs, with AMDV being an outlier.

These findings indicate that the current classification of viruses within *Parvovirinae* (62) does not always reflect their evolutionary relationships. The first discrepancy was found for avian parvoviruses, now classified as members of the *Parvovirus* genus but revealed by our analysis to be linked evolutionarily to AAV rather than to any autonomous parvoviruses (Fig. 1 and 2). Our results concur with an observation on the relatively high homology between GPV, MDPV, and AAV-2 (18, 68) but do not show that GPV is even closer to AAV-1 and AAV-3, an earlier notion based on DNA cross-hybridization data (18). We found that all serotypes of AAV are equidistant from each of the three avian parvoviruses (Fig. 2).

Other discrepancies were found for the simian parvovirus from the long-tailed macaque, ChipmunkPV, and BovinePV. While these three viruses are classified as members of the *Parvovirus* genus, their evolutionary linkage to all known autonomous primate parvoviruses, and not to any known nonprimate parvoviruses, was revealed in our study (Fig. 1 and 3). Our observations concur with recent data on the genetic homology of primate (27) and chipmunk (67) parvoviruses to B19.

A reliable analysis of phylogenetic relationships among the three identified groups of *Parvovirinae* (Fig. 1) was obstructed by the high evolutionary distances. The topology of the phylogenetic tree, in which all three groups of *Parvovirinae* branch out from basically a single phylogenetic node, is likely to reflect the saturation of nucleotide substitutions among the groups. In contrast to intergroup relationships, intragroup relationships could be analyzed in detail.

Another important issue of parvovirus classification is related to the recognition of individual virus species. For several other viruses, such as HIV type 1 (HIV-1), genetic distances among isolates can be higher than 0.3 (38, 39, 41) and biological and immunological characteristics of virus isolates are highly variable (for review, see reference 40). Nevertheless, all HIV-1 strains are considered to belong to the same virus species based on their common evolutionary origin. This principle is used for some parvoviruses but not for others. For example, all five available full-length genome sequences of MVM are considered to be derived from a single species, while the three available genome sequences of MousePV are classified as belonging to three different species, MousePV-1, -1b, and -1c, even though genetic heterogeneity in MVM is actually much higher than in MousePV (Fig. 4). Similarly, the genetic distances among AMDV isolates, which are considered to belong to a single species, are not different from or even higher than those between the two duck parvoviruses or between AAV-3 and -3B or among parvoviruses from carnivores: FelinePV, CaninePV, RaccoonPV, and MEV. The parvoviruses of carnivores have been considered as (host range) variants of a single virus species (46), and our data suggest similar consideration for MousePV-1, -1b, and -1c, the two duck parvoviruses, AAV-3 and -3b, and possibly AAV-1 and -6.

Driving forces of parvovirus evolution. To study evolutionary forces that are operational among parvoviruses, we analyzed synonymous versus nonsynonymous nucleotide substitutions in the two ORFs. Nonsynonymous substitutions, as they

change the amino acids, are generally subjected to strong positive or negative selection pressure. In contrast, synonymous substitutions, which preserve amino acids, are supposed to be subjected to a weaker selection pressure or to none. Since the mutation rates at synonymous and nonsynonymous sites should be the same, D_s and D_a , as well as their ratios, indicate the direction and intensity of selection in the evolutionary history of a group of species. For instance, the D_s/D_a ratios among HIV-1 polymerase sequences are well above 1, since most nonsynonymous substitutions within this gene are deleterious (23). In contrast, short-term intrahost evolution of the HIV-1 envelope gene is characterized by mean D_s/D_a ratios of 0.4, reflecting the advantageous character of amino acid changes in this immunogenic region (41). For long-term evolution, as the separation among HIV-1 subtypes, the D_s/D_a ratios within the *env* gene are generally above 1 (39), reflecting accumulation of synonymous substitutions with time (26).

For all pairwise sequence comparisons, except for the extremely homogeneous PorcinePV, we found D_s/D_a ratios above 1. The most extreme case of negative selection was observed for the separation between the B19 and V9 lineages, apparently an ancient event. While these two viruses were extremely homologous at the amino acid level, with the mean D_a between them not exceeding 0.03, the mean D_s were 0.40 to 0.45 resulting in D_s/D_a ratios of up to 22.5 (Fig. 3). The D_s/D_a ratios were above 1 even for recent evolutionary events, such as the cross-species transmission of FelinePV to dogs (35, 46, 49), when an increase of nonsynonymous substitutions during virus adaptation to a new host could be expected. We did observe a two- to four-times-lower genetic heterogeneity among CaninePV than among FelinePV, a likely indication of a recent transmission bottleneck. At the same time, the mean D_s/D_a ratios were 5.8 for orf1 and 3.3 for orf2 for the comparisons of CaninePV to FelinePV. Among CaninePV, the mean D_s/D_a ratios were 8.0 for orf1 and 2.0 for orf2, compared to a mean ratio of 4.0 for both ORFs among FelinePV. While these data do not exclude the possibility that certain nonsynonymous substitutions were selected for during the adaptation of FelinePV to dogs (35), they indicate that the influence of positive selection during this recent cross-species transmission was extremely limited.

In contrast to that for the cross-species transmissions of FelinePV to dogs (46), the time scales for separation between and diversification within other virus species are not known. Since little is known about the evolution rate of parvoviruses, precise dating of those events is currently not possible. CaninePV has been shown to evolve in a linear fashion over time with the mean evolution rate of 10^{-4} nucleotides per year (35), which would require 100 years of independent evolution for the evolutionary distance of 0.01 between two lineages. Apparently, this evolution rate has to be considered maximal, since it is measured during a short period of virus adaptation to a new host. Moreover, we demonstrated that the evolution rate of parvoviruses is far from being uniform for synonymous and nonsynonymous positions.

Our analysis provided evidence for both host-dependent and independent evolution in the history of parvoviruses. Within two phylogenetic groups, the autonomous primate parvoviruses and AAV and avian parvoviruses, the phylogenetic relationships were host dependent. For instance, the relationships among B19, V9, and parvoviruses from three macaque species

matched exactly the relationships among their hosts, according to an earlier analysis of primate mitochondrial DNA (30). On the other hand, human B19 and V9 viruses and AAV were evolutionarily related to simian and avian parvoviruses, respectively, rather than to each other. For the third phylogenetic group, in the homogeneous subgroup of viruses from carnivores and, most remarkably, the heterogeneous subgroup of rodent viruses, no host-specific clusters were observed (Fig. 4). While, unlike what was found for viruses from carnivores (46), there is no epidemiological evidence for cross-species transmissions of rodent parvoviruses, many of them are able to experimentally infect different hosts. For instance, LuIII can establish infection in hamsters (59). The absence of host-specific clusters and genetic mosaicism of rodent parvoviruses suggest that cross-species transmissions may have occurred among rodents.

For most virus comparisons, the topologies of phylogenetic trees were similar in both ORFs, with two exceptions. First, the positions of AAV-2 and AAV-4 in the phylogenetic trees varied in relation to the genomic region analyzed (Fig. 2B to D versus E to G). Taken with our observations of pairwise D_a among various viruses being drastically higher or lower within orf1 than within orf2, this finding suggests that the selection pressure on the two genomic regions differs among distinct lineages, which could be related to functional difference between the two ORFs.

The second case of tree incongruity was observed among parvoviruses from mice and hamsters. MousePV and HamsterPV were evolutionary related to MVM within orf1 and to LuIII within orf2 (Fig. 4). Since this incongruity was observed for both nonsynonymous and synonymous substitutions, it is unlikely to be the result of convergent evolution. We demonstrated that this genomic mosaicism is likely to be the result of a recombination that occurred among lineages within this group (Fig. 5). Traditionally, MousePV and HamsterPV should be considered the recombinants between MVM and LuIII-related viruses. However, one cannot exclude the possibility that the recombination event involved a yet-undiscovered parvovirus.

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